Bibliographic Fields

Document Identity

(19)□□□□□	(19) [Publication Office]
000000000	Japan Patent Office (JP)
(12)00000	(12) [Kind of Document]
0000000	Japanese Patent Publication (B2)
(11)000000	(11) [Patent Number]
00000000000000000000000000000000000000	Patent No. 3421202*(P3421202)
(45)	(45) [Issue Date]
00000000000000000000000000000000000000	Heisei 15*June 30 days (2003.6.30)
(43)□□□□	(43) [Publication Date of Unexamined Application]
00000000000000000000000000000000000000	1998 April 28*(1998.4.28)
Filing	
(24)□□□□□	(24) [Registration Date]
00000000000000000000000000000000000000	Heisei 15*April 18*(2003.4.18)
(21)	(21) [Application Number]
0000000000	Japan Patent Application Hei 8- 268385
(22)□□□□□	(22) [Application Date]
000000000000000000000000000000000000000	1996 October 9*(1996.10.9)
000000	*Request for Examination **
000000000000000000	2000 April 4*(2000.4.4)
Public Availability	•
(45)□□□□□	(45) [Issue Date]
00000000000000000000000000000000000000	Heisei 15*June 30 days (2003.6.30)
(43)□□□□	(43) [Publication Date of Unexamined Application]
00000000000000000000000000000000000000	1998 April 28*(1998.4.28)
Technical	
(54)	(54) [Title of Invention]
00000000000000000000000000000000000000	PROPYLENE POLYMER WHICH IS ACQUIRED MAKING USE OF POLYMERIZATION METHOD OF PROPYLENE ANDTHAT
(51)00000000700	(51) [International Patent Classification, 7th Edition]

C08F 2/02	C08F 2/02
4/654	4/654
10/00 510	10/00 510
□FI□	[FI]
C08F 2/02	C08F 2/02
4/654	4/654
10/00 510	10/00 510
000000	[Number of Claims]
	8
abaaa	[Number of Pages in Document]
	13
(56)	(56) [Cited Reference(s)]
0000	[Literature]
00 00000000000000	Japan Unexamined Patent Publication Sho 64- 69610(JP,A)
	[Literature]
00 000000000000000	Japan Unexamined Patent Publication Hei 7- 31 6220(JP,A)
	[Literature]
00 00000000000000	Japan Unexamined Patent Publication Hei 7- 179514(JP,A)
	[Literature]
00 000000000000	Japan Unexamined Patent Publication Hei 8- 231 625(JP,A)
	[Literature]
00 000000000000	Japan Unexamined Patent Publication Hei 9- 52911(JP,A)
(58)	(58) [Field of Search]
□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	(International Class 7,DB*)C08F 2/00 - 2/60,4/654
(65)	(65) [Publication Number of Unexamined Application (A)]
0000000000	Japan Unexamined Patent Publication Hei 10- 110 003
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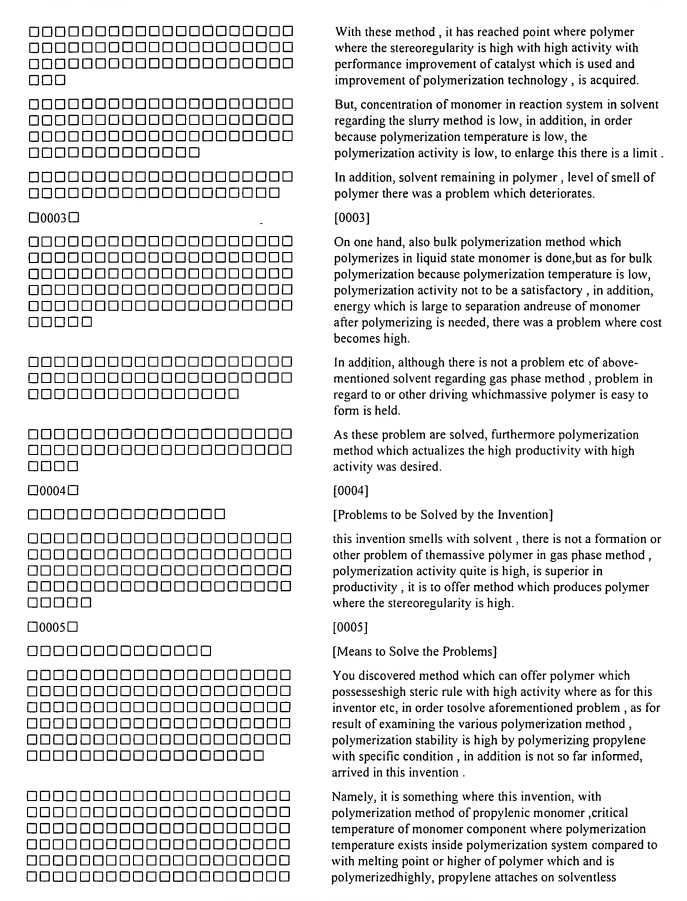
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Claims	
(57)	(57)[Claim(s)]
000010	[Claim 1]
00000000000000000000000000000000000000	Other than propylene, the;al to ethylene or carbon number 4 or more 12 0.01 - 30 weight % it is possible to include -olefin as comonomer, with polymerization method of the propylenic monomer, critical temperature of monomer component where polymerization temperature exists inside polymerization system compared to with melting point or higher of polymer which and is polymerizedhighly, it is higher than critical pressure of monomer component where polymerization pressure exists inside aforementioned polymerization system, polymerization temperature 170 deg C~250 deg C, polymerization pressure is inside range of 50 - 300 kg/cm ² , propylene itattaches on solventless polymerization under condition and makes feature, the

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	polymerization method . of aforementioned propylene
000020	[Claim 2]
00000000000000000000000000000000000000	Q of polymer is 6 or more , polymerization method . of propylene which isstated in Claim 1
00030	[Claim 3]
0000000000000001000	polymerization method . of propylene which uses Ziegler catalyst , states in Claim 1
0000 40	[Claim 4]
00000000000000000000000000000000000000	polymerization method . of propylene which uses magnesium- bearing catalyst , states in Claim 1
	[Claim 5]
000000100000000000 0000001000000000000	polymerization method . of propylene which uses catalyst which prepolymerization is donewith vinyl group-containing compound , states in Claim 1
	[Claim 6]
00000000000000000000000000000000000000	melting point of polymer which prepolymerization is done is $200 \ deg \ C$ or greater , polymerization method . of propylene which is stated in Claim 5
000070	[Claim 7]
00010000000000000000000000000000000000	With polymer which was polymerized with method which is stated in Claim 1, MFR of said polymer is $5 - 1000 \text{g/}10 \text{min}$, it makesfeature, propylene polymer .
000080	[Claim 8]
00010000000000000000000000000000000000	With polymer which was polymerized with method which is stated in Claim 1, Q of said polymer is 6 or more, it makesfeature, propylene polymer.
Specification	
00000000	[Description of the Invention]
□0001□	[0001]
00000000000	[Technological Field of Invention]
00000000000000000000000	this invention is something regarding polymerization method of propylene .
000000000000000000000000000000000000000	Furthermore details regard polymerization method of propylene which possesses the high stereoregularity with quite high activity.
□0002□	[0002]
000000	[Prior Art]
	Until recently as method which polymerizes propylene, uses the hexane, heptane etc as solvent gas phase method etc which designates bulk method, inert gas whichdesignates slurry method, propylene itself which as solvent as media is known.

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JP3421202B2 2003-6-30 polymerization under condition which ishigher than critical 0000000000000000 pressure of monomer component where polymerization pressure exists inside theaforementioned polymerization system makes feature, offers polymerization method of theaforementioned propylene. □0006□ [0006] 0000000000 [Embodiment of the Invention] 00000000000000000000 this invention with temperature which is higher than critical 0000000000000000000 temperature of monomer component which exists inside 00000000000000000 polymerization system and is polymerized with temperature of the melting point or higher of polymer which is polymerized. Namely, it makes temperature range of melting point or higher of polymer where polymerization temperature when from critical temperature of propylene which is offered to thepolymerization, in addition producing copolymer, is high 0040000in comparison with all critical temperature of propylene and other comonomer component (You detail on description below, but the;al of ethylene and carbon number 4 or more -olefin etc) which are used, atsame time forms. □0007□ [0007] In addition, polymerization pressure makes high pressure in comparison with the critical pressure of monomer component which exists inside polymerization system. 00000000000000000000 Furthermore in this invention, "melting point of polymer" with, polymerizing under same condition as condition which is done beforehand actually, as a result it meansthat melting point of produced polymer which is acquired was measured due to the differential thermal analysis (DSC). □0008□ [0008] When polymerization temperature, it is lower than critical temperature of monomer component which existsinside polymerization system, monomer for polymerization doing. liquefaction because density of monomer becomes high, as benefit of this invention becomes impossible is enjoyed, cost of separation and reuse of monomer becomes high. 00000000000000000000 Furthermore, when being lower than melting point of polymer , molecular weight distribution which is made objective of QDD6DDDDDDDDDDDDDDDDD this invention it is wide, preferably Q stops being acquired, propylene polymer of 6 or more is not desirable.

[0009]

□0009□

In addition, when polymerization pressure, being lower than critical pressure of monomer component which exists inside polymerization system, supercritical phenomenon cannot reveal and cannotacquire high activity.

As for polymerization temperature and polymerization pressure homopolymerization or copolymerization?, or



000-000-1000000000	
00000000000000000000000000000000000000	Quantity of comonomer is not something which especially isrestricted. They are 15 weight %extent from 0.01 - 30 weight %, preferably 0.1.
□0013□	[0013]
□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	In addition, following to method of this invention, as for propylene polymer which was polymerized, it is desirable for MFR to be 5 - 1000 g/10 min
□□□□MFR□ASTMD1238□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	Furthermore, this MFR conforming to ASTM D1238, is something which ismeasured, (230 deg C, 2.16kg load).
□0014□	[0014]
00000000000000000000000000000000000000	Furthermore, following to method of this invention, as for propylene polymer which was polymerized, it is desirable for Q to be 6 or more.
00000000000000000000000000000000000000	When Q is this range, moldability of mechanical strength corresponding and external appearance of molded article are superior, when it is too low, there is a tendency where external appearance of moldability and molded article is inferior.
	Furthermore Q is something which is calculated with Mw/Mn (weight average molecular weight, Mn which was measured by Mw:gel permeation chromatograph (GPC) is number average molecular weight which wasmeasured by GPC.).
□0015□	[0015]
00000000000000000000000000000000000000	As for catalyst which is used with this invention, it can recognize theeffect of this invention, if you obtain with those of option, butbecause Ziegler catalyst, especially being a catalyst of magnesium bearing type is superior, activity in high temperature, it is desirable.
00000000000(A)0000(B)00 0000(A)000(B)0000(C)0000 0000000000	Concretely, below-mentioned component (A) and component (B), or component (A), component (B) and it is something which consists of combination of component (C).
00000000000000000000000000000000000000	It is not something where here, "Of combination it consists" with as for notion that whereyou say, that component which is used is only raising/holdingShimesu thing (namely, component (A) and (B), component (A), (B) and (C)) you say and mean, coexistence of goal-consistent other component is not removed.
□0016□	[0016]
00(A)000000000000000000 0000000000000000	<component a="">component (A) is solid component for Ziegler catalyst which contains titanium, magnesium and halogen as essential ingredient.</component>
00000000000000000000000000000000000000	Here "It contains as essential ingredient" with may include goal-consistent other element outside raising/holding Shimesu three components to say these element each one may exist as

	compound of the goal-consistent option, and these element may exist as is connected mutually, it issomething which is shown.
00000000000000000000000000000000000000	solid component itself which includes titanium, magnesium and halogen is somethingof public knowledge.
	for example Japan Unexamined Patent Publication Showa 53-45688 number, same 54 - 3894, same 54 - 31,092, same 54 - 39483, same54 - 94591, same 54 - 118484, same 54 - 131,589, same 55 - 75411, same55 - 90510, same 55 - 90511, same 55 - 127405, same 55 - 147507, same55 - 155003, same 56 - 18609, same 56 - 70005, same 56 - 72001, same56 - 86905, Same 56 - 90807, same 56 - 155206, same 57 - 3803, same 57 - 34103, same 57 - 92007, same 57 - 121003, same 58 - 5309, same 58 - 5,310, same 58 - 5,311, same 58 - 8706, same 58 - 27732, same 58 - 32604, same 58 - 32605, same 58 - 67703, same 58 - 117206, same 58 - 127708, same 58 - 183708, Same 58 - 183709, those which are stated in same 59 - 149905, same 59 - 149906 each disclosure etc are used.
□0017□	[0017]
	Regarding to this invention, you can list carbonate etc of magnesium halide, di alkoxy magnesium, alkoxy magnesium halide, magnesium oxyhalide, dialkyl magnesium, magnesium oxide, magnesium hydroxide, magnesium as the magnesium compound which becomes magnesium source which is used.
0000000000000000000 000000000000000000	Desirable ones are magnesium halide, di alkoxy magnesium, alkoxy magnesium halide among these.
□0018□	[0018]
00000000000000000000000000000000000000	In addition, as for titanium compound which becomes titanium source, it can increase compound which is displayed with General Formula Ti (OR') _{4- q} X _q (As for R' with hydrocarbon residue, with those of preferably carbon number $1\sim10$ extent, as for the X halogen is shown here, q 0 <= shows quantity of $q*4$.).
0000000	As embodiment,
TiCl₄ □	TiCl ₄ ,
TiBr₄ □	TiBr ₄ .

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 $Ti(OC_2 H_5)Cl_3 \square$ $Ti(OC_2 H_5)_2 Cl_2 \square$

□0019□

000TiXD₄ (0000X0000000000)00000 00000000

□0020□

Ti (OC₂ H₅) Cl₃ ,

Ti (OC₂ H₅) ₂ Cl₂ ,

You can list Ti (OC₂ H₅) ₃ Cl, Ti (O-iC₃ H₇) Cl₃, Ti (O-nC₄ H₉) $Cl \leq b \leq 3 \leq sub \leq 1$, $Ti (O-nC \leq sub \leq 4 \leq sub \leq H \leq sub \leq 9 \leq sub \leq 1$ ₂ Cl₂, Ti (OC₂ H₅) Br₃, Ti (OC₂ H₅) (OC₄ H₉) ₂ Cl, Ti (O-nC₄ H₉) ₃ Cl, Ti (O-C₆ H₅) C1 < sub > 3 < /sub >, Ti (O-iC < sub > 4 < /sub > H < sub > 9 < /sub >)₂ Cl₂ , Ti (OC₅ H₁₁) C₁₃ , Ti (OC₆ H₁₃) C₁₃, Ti (OC₂H₅) ₄, Ti (OnC₃ H₇) ₄ , Ti (OnC₄ H₉) ₄ , Ti (OiC₄ H₉) ₄ , Ti (OnC₆ H₁₃) ₄ , Ti (OnC₈ H₁₇) ₄ , Ti {OCH₂ CH (C₂ H₅) $C \le b \le 4 \le b \le 4 \le b \le 4 \le b \le 4 \le b \le etc.$

[0019]

In addition, electron donor which it mentions later in TiX*₄ (Here as for X* halogen is shown.) the molecular compound which reacts can be used.

As embodiment, you can list TiCl₄ *
CH₃ COC₂ H₅,
TiCl₄ * CH₃ CO₂
C₄ * CH₃ CO₂
C₆ H₅,
TiCl₄ *
C₆ H₅ NO₂,
TiCl₄ *
CH₃ COCl,
TiCl₄ *
C₆ H₅ H₅ COCl,
TiCl₄ *
C₆ H₅ H₅
CO₂ H₅
CO₂ H₅,
TiCl₄ *
CO₂ H₅ H₅,
TiCl₄ *
CICOC₂ H₅ H₅ ,
TiCl₄ *
CC₄ H₅ H₅ ,
TiCl₄ *
CC₄ H₄ O etc.

Even in these titanium compound desirable ones are TiCl₄, Ti (OC₂ H₅) ₄, Ti (OC₄ H₉) ₄, Ti (OC₄ H₉) Cl₃ etc.

[0020]

As halogen source, it is normal to be supplied from halogen compound of theabove-mentioned magnesium and/or titanium, but it is possible also to supply from the halogenating agent of public knowledge such as halide of aluminum and halide of

JP3421202B2 2003-6-30 the halide, phosphorus of silicon. 0000000000000 halogen which is included in catalyst component fluorine, chlorine, bromine, iodine or is good even with these blend, especially chlorine is desirable. □0021□ [0021] As for solid component which is used for this invention, also use of SiCl₄, CH₃ SiCla SiCl₃ or other silicon compound, methyl hydrogen polysiloxane or other polymeric silicon compound, $(OiC_3 H_7)_3 \square AlCl_3 \square AlBr_3 \square Al(OC_2 H_5)_3$ Al (OiC₃ H₇)₃ , Al □AI(OCH₃ Cl₃, Al Br₃, Al (OC < sub > 2 < /sub > H < sub > 5 < /sub >) < sub > 3 < /sub > , Al)3 \square B(OC₂ H_5 \square B(OC₆)3 (OCH₃) ₂ Clor other aluminum)3 ☐ MoCl₅ compound and B (OCH₃) ₃, B 0000000000000000000000 (OC₂ H₅) ₃ , B (OC < sub > 6 < / sub > H < sub > 5 < / sub >) < sub > 3 < / sub > or otherborated compound, WCI₆, MoCI₅ or other other component being possible to other than abovementioned essential ingredient, these as forremaining in solid component as silicon, aluminum and boron or other component there is not aninconvenience. Furthermore, when this solid component is produced, using electron donor as the internal donor, it is possible also to produce. □0022□ [0022] esters, ethers, diether of alcohols, phenols, ketones, aldehydes, carboxylic acid and organic acid or inorganic acid and nitrogen-containing electron donor etc like oxygen 0000000000000000000 containing electron donor, ammonia, amine, nitrile, isocyanate like acyl amides, acid anhydride are illustrated is possible electron donor which can be utilized in production of this solid component (internal donor) as. □0023□ [0023] More concrete, (0)000000(J2) methanol, ethanol, propanol, pentanol, hexanol,

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octanol.

dodecanol,

octadecyl alcohol,

000000



00000000000000000000000000000000000000	can be listed.
00000000000000000000000000000000000000	2 kinds or more you can use these electron donor.
00000000000000000000000000000000000000	As for being desirable among these with organic acid ester and organic acid halide, and diether, as for especially being desirable it is a phthalic acid ester, cellosolve acetate, phthalic acid halide and diether.
□0024□	[0024]
00000000000000000000000000000000000000	As for amount used of above-mentioned each component, it can recognize effect of this invention, if it can with those of option, butgenerally, inside of following range is desirable.
□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	As for amount used of titanium compound, inside of range of 1 X 10 ⁻⁴ -1000 is goodwith mole ratio vis-a-vis amount used of magnesium compound which is used, it is inside range of preferably 0.01~10.
00000000000000000000000000000000000000	When compound for that is used as halogen source, amount used titanium compound and/or magnesium compound includes halogen, it is inside range of 1 X 10 ⁻² ~1000, preferably 0.1 ~100, with mole ratio vis-a-vis amount used of magnesium which is used it does not include ofregardless.
□0025□	[0025]
00000000000000000000000000000000000000	amount used of silicon, aluminum and borated compound is inside range of 1 X 10 ⁻³ ~100, preferably 0.01~1, with mole ratio vis-a-vis amount used of above-mentioned magnesium compound.
00000000000000000000000000000000000000	amount used of electron donating compound is inside range of 1 X 10 ⁻³ ~10, preferably 0.01~5, with mole ratio vis-a-vis amount used of above-mentioned magnesium compound.

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□0026□	[0026]
00(A)0000000000000000000000000000000000	solid component in order to produce component (A) above- mentioned titanium source, magnesium source and halogen source, furthermore is produced making use of electron donor or other other component, likebelow for example by production method in accordance with necessary.
(0)00000000000000000000000000000000000	(J2) magnesium halide and according to need electron donor and titanium-containing compound method . which contacts
(0)00000000000000000000000000000000000	method. which treats (jp2) alumina or magnesia with halogenated phosphorus compound, the magnesium halide, electron donor, titanium halogen containing compound contacts that
(0)00000000000000000000000000000000000	(jp3) magnesium halide and titanium tetraalkoxide and specific polymeric silicon compound contacting, in the solid component which is acquired, halogen compound, and according to need electron donor of titanium halogen compound and/or silicon the method. which contacts
00000000000000000000000000000000000000	As this polymeric silicon compound, those which are shown with formula below are suitable.
□0027□	[0027]
0010	[Chemical Formula 1]
$ \begin{array}{c} $	
□0028□	[0028]
(DDDR ² DDDD1~10DDDDDDDDDDrDDD DDDDDDDDDDDDDDDDDDDDDDD	(Here, as for R ² as for hydrocarbon residue, r of carbon number 1~10extent, kind of degree of polymerization where viscosity of this polymeric silicon compound becomes 1 - 100 centistokes extent is shown.)
00000000000000000000000000000000000000	Among these, methyl hydrogen polysiloxane, 1,3,5,7-tetramethyl cyclotetrasiloxane, 1,3,5,7,9-pentamethyl cyclopentasiloxane, ethyl hydrogen polysiloxane, phenyl hydrogen polysiloxane, cyclohexyl hydrogen polysiloxane etc is desirable.
□0029□	[0029]
(0)00000000000000000 000000000000000000	Melting (jp4) magnesium compound with titanium tetraalkoxide and electron donor, method which the titanium

compound contacts solid component which it precipitated

00000 00000	with halogenating agent or the titanium halogen compound
(0)00000000000000000000000000000000000	(jp5) Grignard reagent or other organomagnesium compound halogenating agent, reductant etc and after action, in this according to need electron donor and the titanium compound method. which contacts
(B)000000000000000000000000000000000000	In (jp6) alkoxy magnesium compound halogenating agent and/or titanium compound method. which contacts under existing or absence of electron donor
□0030□	[0030]
00000(A)00000000000 000000000	As manufacturing condition of component (A) of this invention, inside ofbelow-mentioned range is desirable.
□□□□□□-50~200 deg □□□□□□□□□0~100 deg □□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	contact temperature - is 50 - 200 deg Cextent , preferably 0~100 deg Cextent , .
	As contact method, mechanical method with such as rotating ball mill, vibrating mill, jet mill, medium-stirring mill. Under existing of inactivity diluent, you can list method etc which contacts with churning.
00000000000000000000000000000000000000	You can list hydrocarbon and halohydrocarbon, polysiloxane etc of aliphatic or aromatic as inactivity diluent which this time is used.
□0031□	[0031]
0000000000(A)00000000 000000000000000000	component (A) which is used with this invention can also use as those whichpass prepolymerization step which consists of fact that contacting, youpolymerize vinyl group-containing compound, for example olefins, diene compound, styrene etc.
00000000000000000000000000000000000000	As embodiment of olefins which is used occasion where prepolymerization is done, those of for example carbon number 2~20extent . There is a ethylene , propylene , 1-butene , 3- methyl butene -1,1- pentene , 1- hexene , 4- methylpentene -1,1- octene , 1- decene , 1- undecene , 1-eicosene etc concretely, there are 1 and 3-butadiene , isoprene , 1,4- hexadiene , 1,5-hexadiene , 1,3- pentadiene , 1,4- pentadiene , 2,4- pentadiene , 2,6-octadiene , cis-2,trans -4- hexadiene , trans -2, trans -4- hexadiene , 1,3- heptadiene , 1,4- heptadiene , 1,5-heptadiene , 1,6-heptadiene , 2,4- heptadiene , dicyclopentadiene , 1,3- cyclohexadiene , 1,4- cyclohexadiene , cyclopentadiene , 1,3- cyclo heptadiene , 4- methyl -1,4- hexadiene , 5-methyl -1,4- hexadiene , 1,9- decadiene , 1,13- tetradecadiene , p- divinyl benzene , m- divinyl benzene , o-divinyl benzene , dicyclopentadiene etc as the embodiment of diene compound .

1,3-000000000004-000-1,4- 00000001,9-0000001,13- 00000000000-000000000m- 0000000000000	
00000000000000000000000000000000000000	In addition, you can list styrene, ;al-methylstyrene, allyl benzene, chlorostyrene etc as embodiment of the styrene.
□0032□	[0032]
00000000000000000000000000000000000000	As for reaction condition of titanium component and above- mentioned vinyl group-containing compound, it canrecognize effect of this invention, if it can with those of option, but inside of following range is desirable generally.
00000000000000000000000000000000000000	amount of prepolymerization of vinyl group-containing compound per titanium solid component 1 gram 0.001 - 1000 gram, preferably 1~100gram, furthermore is inside range of preferably 5~50gram.
□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	reaction temperature at time of prepolymerization - is 150 - 85 deg C, preferably 0~50 deg C.
00000000000000000000000000000000000000	Low polymerization temperature is desirable and, "This polymerization", namely the; al -olefin in comparison with polymerization temperature when polymerizing.
00000000000000000000000000000000000000	Reaction does generally under agitating, it is desirable, thattime n- hexane, n-heptane or other inert solvent to be possible also, it is possible also, in additionwith types of vinyl compound to do that itself of above-mentioned vinyl compound to exist, as media it is possible also to execute with the gas state.
0000000000000200000 00000000	In addition 2 kinds or more also to jointly use this vinyl group-containing compound it is possible.
00000000000000000000000000000000000000	In addition, as for melting point of polymer which prepolymerization is done, it is desirable from reason of activity improvement of component (A) to be 200 deg C or greater.
□0033□	[0033]
00000(8)0000000000000 000000000000000000	As for component (B) of <component (b)="">this invention, blend, of organo-aluminum compound, or organo-aluminum compound which is displayed with below-mentioned General Formula [II] and isdisplayed with below-mentioned General Formula [III] organo-aluminum compound which are displayed with below-mentioned General Formula [1]</component>
$AIR^3_{3-n}X^1_n\square I\square$	AIR ³ _{3- n} X ¹ _n [1]

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)₂ Al Cl, (n- C₄ H₉)₂ Al Cl, (i- C₄ H₉)
Al Cl₂ , (n- C₆ H₁₃)
₂ Al Cl, (n- C₈ H₁₇)
₂ Al Cl, (n- C₁₀ H₂₁)

₂ Al Cl and a (C₂ H₅)₂ Al Br etc as embodiment of organo-aluminum

	compound which is displayed with General Formula [II].
□0036□	[0036]
0000111 0000000 000000	As embodiment of organo-aluminum compound which is displayed with General Formula [III],
$(CH_3)_2 Al(OC_2 H_5) \square$	(CH ₃) ₂ Al (OC ₂ H ₅),
$(C_2 H_5)_2 Al(OCH_3) \square (C_2 H_5)_2 Al(OC_2 H_5) \square (i-C_3 H_7)_2 Al(OC_2 H_5) \square (n-C_6 H_{13})_2 Al(OC_4 H_9)_2 Al(OC_2 H_5) \square (n-C_6 H_{13})_2 Al(OC_4 H_9) \square (n-C_8 H_{17})_2 Al(OCH_3) \square (n-C_{10} H_{21})_2 Al(OC_2 H_5) \square (CH_3) Al(OCH_3)_2 \square (C_2 H_5) Al(OC_2 H_5)_2 \square (i-C_3 H_7) Al(OC_4 H_9)_2 \square (n-C_4 H_9) Al(OC_6 H_5)_2 \square (n-C_6 H_{13}) Al(OC_6 H_{13})_2 \square \square \square (n-C_{10} H_{21}) Al(OCH_3)_2 \square \square \square \square \square$	(C ₂ H ₅) ₂ Al (OCH ₃), (C ₂ H ₅) ₂ Al (OC ₂ H ₅), (i- C ₃ Al (OC ₂ H ₅ Al (OC ₂ Al (OC ₂ Al (OC ₂ H ₅), (n- C ₃ H ₇) ₂ Al (OCH ₃), (n- C ₄ H ₉) (n- C ₃), (n- C ₄ H ₉), (n- C ₆ H ₅), (n- C ₆ H ₁₃) (n- C ₈ H ₁₇) (n- C ₈) (n- C ₁₀ H ₂₁ Al (OCH ₃), (n- C ₁₀ H ₅), there is a (CH ₃) Al (OCH ₃) (C ₂) (C ₂ H ₅) Al (OCC ₂) H ₅) Al (OC ₂) Al (OC ₆) Al (OC ₉) Al (OC ₆) Al (OC ₉) Al (OC ₆) Al (OC ₆) H ₅) Al (OC ₆) H ₁₃) Al (OCH ₃)
□0037□	[0037]
00(B)00000011000000000000000000000000000	When blend of organo-aluminum compound of General Formula [II] and organo-aluminum compound of General Formula [III] is used component (B) as, those mixing ratio, $0.1 - 100$, are inside rangeof preferably $0.1 \sim 10$, with mole ratio of the latter for former.
□□(B)□□□□□□□□(A)□□□□□□□ □□□□□□□□□(Al/Ti)□0.1~1000□□□ □□1~200□□□□□□□□	amount used of component (B) with mole ratio for titanium component in component (A) is inside range of 0.1 - 1000 preferably 1~200 with (Al/Ti).
□0038□	[0038]
0000000000(c)00000000 000000000000000000	Regarding to <component (c)="">this invention, uses electron donor component (C) as to be possible, preferably concretely esters, diether of organic acid or inorganic acid, you can use those which are chosen from ethers and amines.</component>

□0039□	[0039]
	As organic acid esters, 1 - dibasic carboxylic acid of carbon number 1~20 or 1 - dihydric alcohol of carbon number 1~20 of carbon dioxide (With this invention, it handles carbon dioxide as organic acid) (ether alcohol is included.) with ester (intramolecular ester is included.) is representative.
	methyl formate, methyl acetate, ethylacetate, vinyl acetate, propyl acetate, octyl acetate, cyclohexyl acetate, cellosolve acetate, ethyl propionate, methyl butyrate, ethyl valerate, ethyl stearate, methyl chloroacetate, ethyl dichloroacetate, methyl methacrylate, ethyl crotonate, ethyl cyclohexane carboxylate, methyl benzoate, ethyl benzoate, propyl benzoate, butyl toluate, octyl benzoate, cyclohexyl benzoate, phenyl benzoate, benzyl benzoate, cellosolve benzoate, methyl toluate, ethyl toluate, amyl toluate, ethyl ethylbenzoate, methyl anisate, ethyl anisate, ethyl ethoxybenzoate, diethyl phthalate, dibutyl phthalate, diheptyl phthalate, ;ga-butyrolactone, you can list the;al-valerolactone, coumarin, phthalide, ethylene carbonate etc concretely.
□0040□	[0040]
00000000000000000000000000000000000000	As inorganic acid ester, before, kind of alcohol which was inscribed it canincrease ester silicon, boron, phosphorus and aluminum, or other oxyacid and concerning organic acid ester.
00000000000000000000000000000000000000	Portion of atomic valency of these element may be sufficed hydrocarbon residue (carbon number 1~8extent) orwith halogen atom .
00000000000000000000000000000000000000	In this kind of inorganic acid ester, ester of oxyacid of silicon isdesirable.
□0041□	[0041]
0000000	As embodiment,
(CH ₃)Si(OCH ₃) ₃ □	(CH ₃) Si (OCH ₃) ₃ ,
$(CH_3)Si(OC_2 H_5)_3 \square$	(CH ₃) Si (OC ₂ H ₅) ₃ ,
$(C_2 H_5)_2 Si(OCH_3)_2 \square$	(C ₂ H ₅) ₂ Si (OCH ₃) ₂ ,
$(n-C_6 H_{13})Si(OCH_3)_3 \square$	(n- C ₆ H ₁₃) Si (OCH ₃) ₃ ,
$(C_2 H_5)Si(OC_2 H_5)_3 \square$	(C ₂ H ₅) Si (OC ₂

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$(n-C_{10}H_{21})Si(OC_2 H_5)_3$	(n- C ₁₀ H ₂₁) Si (OC ₂ H ₅) ₃ ,
$(CH_2 = CH)Si(OCH_3)_3$	(CH ₂ =CH) Si (OCH ₃) ₃ ,
Cl(CH ₂) ₃ Si(OCH ₃) ₃ □	Cl (CH ₂) ₃ Si (OCH ₃) ₃ ,
Si(OCH₃)₄□	Si (OCH ₃) ₄ ,
Si(OC ₂ H ₅) ₃ Cl□	Si (OC ₂ H ₅) ₃ Cl,
$(C_2 H_5)_2 Si(OC_2 H_5)_2 \square$	(C ₂ H ₅) ₂ Si (OC ₂ H ₅) ₂ ,
$(C_{17}H_{35})Si(OCH_3)_3$	(C ₁₇ H ₃₅) Si (OCH ₃) ₃ ,
Si(OC ₂ H ₅) ₄ □	Si (OC ₂ H ₅) ₄ ,
$(C_6 H_5)Si(OCH_3)_3 \square$	(C ₆ H ₅) Si (OCH ₃) ₃ ,
Si(OCH ₃) ₂ Cl ₂ □	Si (OCH < sub > 3 < /sub >) < sub > 2 < /sub > Cl < sub > 2 < /sub > ,
$(C_6 H_5)_2 Si(OCH_3)_2 \square$	(C ₆ H ₅) ₂ Si (OCH ₃) ₂ ,
$(C_6 H_5)(CH_3)Si(OCH_3)_2 \square$	(C ₆ H ₅) (CH ₃) Si (OCH ₃) ₂ ,
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	(C ₆ H ₅) Si (OC ₂ H ₅) (i- C ₃ H ₅) (i- C ₃ H ₇) (i- C ₃ H ₇) Si (OCH ₃) Si (OCH ₃) Si (OC ₂) Si (OC ₂) Si (OC ₂ H ₅) Si (OC ₄ H ₅) Si (OCH ₃) (i- C ₄ H ₉) Si (OCH ₃) (C ₆ Si (OCH ₃) Si (OCH ₃) Si (OCH ₃) Si (OCH ₃) (C ₆ H ₁₁) (CH ₃) Si (OCH ₅) Si (OCH ₅) Si (OCH ₅) Si (OCH ₅) Si (OCH ₃) Si (OCH ₅) Si (OC ₂ H ₅) Si (OC ₂ H ₅) Si (OC ₂ H ₇) Si (OC ₃ H ₅) Si (OC ₃) Si (OC ₃ H ₅) Si (OC ₃ H ₅) Si (OC ₃ H ₅) Si (OC ₃) Si (OC <sub< td=""></sub<>
□0042□	[0042]
	[Chemical Formula 2]

□0043□ [0043] (CH₃) ₃ CSi (CH₃) $(CH_3)_3 CSi(CH_3)(OCH_3)_2 \square$ $(OCH \le sub \ge 3 \le sub \ge 2 \le sub \ge 3$ (CH₃) ₃ CSi (HC $(CH_3)_3 CSi(HC(CH_3)_2)(OCH_3)_2 \square$ (CH₃) ₂) (OCH₃) ₂, (CH₃) ₃ CSi (CH₃) $(CH_3)_3 CSi(CH_3)(OC_2 H_5)_2\square$ (OC < sub > 2 < /sub > H < sub > 5 < /sub >) < sub > 2 < /sub > ,(CH₃) ₃ CSi (C₂ $(CH_3)_3 CSi(C_2 H_5)(OCH_3)_2 \square$ H₅) (OCH₃) ₂, (C₂ H₅) ₃ CSi $(C_2 H_5)_3 CSi(CH_3)(OCH_3)_2 \square$ (CH < sub > 3 < /sub >) (OCH < sub > 3 < /sub >) < sub > 2 < /sub > ,(CH₃) ₃ CSi (n- C₃ $(CH_3)_3 CSi(n-C_3 H_8)(OCH_3)_2 \square$ $H \leq sub \geq 8 \leq sub \geq 1$ (OCH $\leq sub \geq 3 \leq sub \geq 1$) $\leq sub \geq 2 \leq sub \geq 1$ $(CH_3)(C_2H_5)CH-Si(CH_3)(OCH_3)_2$ (CH₃) (C₂ H₅) CH-Si (CH₃) (OCH₃) ₂, (CH₃) (CH₃)□ CSi (s -C₃ H₈) $CSi(sec-C_3 H_8)(OCH_3)_3 \square$ (OCH < sub > 3 < /sub >) < sub > 3 < /sub > ,((CH₃) ₂ CHC H₂) $((CH_3)_2 CHCH_2)Si(OCH_3)_2 \square$ $Si(OCH \le sub \ge 3 \le sub \ge 2 \le sub \ge 3$ (CH₃) ₃ CSi (i- $(CH_3)_3 CSi(i-C_3H_8)(OCH_3)_2 \square$ C₃ H₈) (OCH₃) $\leq sub \geq 2 \leq sub > 1$

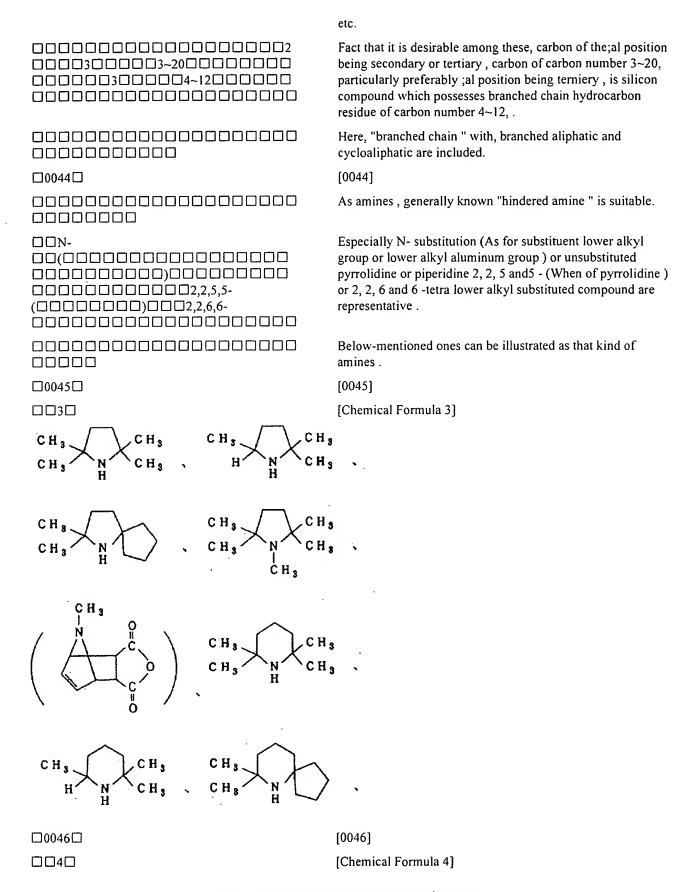
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2003-6-30

C₂ H₅ C (CH₃) C₂ H₅C(CH₃)₂ Si(CH₃)(OCH₃)₂ ₂ Si (CH₃) (OCH₃) ₂. $(CH_3)_3 CSi(n-C_4H_9)(OCH_3)_2 \square$ (CH₃) ₃ CSi (n-C₄H₉) (OCH₃) ₂, C₂H₅C (CH₃) $C_2H_5C(CH_3)_2$ Si(CH₃)(OC₂ H₅)₂ ₂ Si (CH₃) (OC₂ H₅) ₂, (CH₃) ₃ CSi (OCH₃ (CH₃)₃ CSi(OCH₃)₃ □ $) \leq sub \geq 3 \leq sub \geq 3$ $(CH_3)_3 CSi(OC_2 H_5)_3 \square$ (CH₃) ₃ CSi (OC₂ H < sub > 5 < /sub >) < sub > 3 < /sub > ,(C₂ H₅) ₃ CSi $(C_2 H_5)_3 CSi(OC_2 H_5)_3 \square$ (OC < sub > 2 < /sub > H < sub > 5 < /sub >) < sub > 3 < /sub > , $(CH_3)_3 CSi(i-C_4 H_9)(OCH_3)_2 \square$ (CH₃) ₃ CSi (i- C₄ H < sub > 9 < /sub >) (OCH < sub > 3 < /sub >) < sub > 2 < /sub > ,(CH₃) (C₂H₅) CHSi $(CH_3)(C_2H_5)CHSi(OCH_3)_3$ (OCH < sub > 3 < /sub >) < sub > 3 < /sub >(CH₃) ₃CSi (s -C₄ $(CH_3)_3CSi(sec-C_4H_9)(OCH_3)_2$ $H \leq sub \geq 9 \leq sub \geq 1$ (OCH $\leq sub \geq 3 \leq sub \geq 1$) $\leq sub \geq 2 \leq sub \geq 1$ (i- C₃ H₇) ₂ Si (i-C₃ H₇)₂ Si(OCH₃)₂ □ (OCH < sub > 3 < /sub >) < sub > 2 < /sub > , $(C \leq b \leq \leq sub \leq H \leq b \leq 1 \leq sub \leq 3 \leq sub \leq$ $(C_6 H_{11})(CH_3)Si(OCH_3)_2 \square$ (OCH < sub > 3 < /sub >) < sub > 2 < /sub > ,(CH₃) ₃ CSi (n- C₆ $(CH_3)_3 CSi(n-C_6 H_{13})(OCH_3)_2 \square$ $H \leq sub \geq 13 \leq sub \geq 0$ (OCH $\leq sub \geq 3 \leq sub \geq 0$) $\leq sub \geq 2 \leq sub \geq 0$, $(C_6 \ H_{11})_2 \ Si(OCH_3)_2 \ \Box(CH_3)_3 \ CSi(C_6)$ You can list (C₆ H₁₁) $\langle sub \rangle 2 \langle sub \rangle Si (OCH \langle sub \rangle 3 \langle sub \rangle) \langle sub \rangle 2 \langle sub \rangle$ H_{11})(OCH₃)₂ \square (i-C₃ H_7)₂ Si(OC₂ H_5)₂ \square ((CH₃ (CH₃) ₃ CSi (C₆ $_{3}$ C)₂ Si(OCH₃)₂ \square (CH₃)₃ CSi(C₅ H₅)(OCH₃)₂ \square (i-C₄ H₉)₂ Si(OCH₃)₂ \square HC(CH₃)₂C(CH₃)₂ H₁₁) (OCH₃) ₂, (i-C₃ H₇) ₂ Si $Si(CH_3)(OCH_3)_2 \square (C_6 H_{11})(CH_3)Si(OC_2 H_5)_2$ (OC < sub > 2 < /sub > H < sub > 5 < /sub >) < sub > 2 < /sub > , \square HC(CH₃)₂ C(CH₃)₂ Si(OCH₃)₃ \square (C₅H₅)₂ ((CH₃) ₃ C) ₂ Si $Si(OCH_3)_2 \square (i-C_3)$ H_7)Si(OCH₃)₃ (OCH₃) ₂, (CH₃) ₃ CSi (C₅ H₅) (OCH₃) ₂, (i- C₄ H₉) ₂ Si (OCH₃)₂ , HC (CH₃) ₂C (CH < sub > 3 < /sub >) < sub > 2 < /sub > Si (CH < sub > 3 < /sub >)(OCH₃) ₂, (C₆ H₁₁) (CH₃) Si (OC₂ H₅) ₂, HC (CH₃)₂ C (CH₃) ₂ Si (OCH < sub > 3 < /sub >) < sub > 3 < /sub > ,

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(C₅ H₅) ₂ Si (OCH₃) ₂ , (i- C₃ H₇) Si (OCH₃) ₃



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$$\begin{array}{c|c} CH_3 & CH_3 \\ HN & -OC-(C_8H_{18}) - C-O-NH \\ CH_3 & CH_3 \\ \end{array}$$

□0047□	[0047]
00000000000000000000000000000000000000	As ethers, those which it can call hindered ether are suitable.
00000000000000000000000000000000000000	Especially, at least two it possesses lower alkoxy group on carbon atom of the one, this said carbon atom furthermore those which at least one it possesses 6-member ring substituent is representative.
000000 00000	Below-mentioned ones can be illustrated as that kind of ethers .
□0048□	[0048]
	[Chemical Formula 5]

$$\begin{array}{c}
C H_{8} \\
C (O C H_{8})_{2}
\end{array}$$

$$CH_3 \longrightarrow CH_3$$

$$CH_3 \longrightarrow COC_2H_5)_2$$

□0049□

[0049]

[Chemical Formula 6]

□0050□

[0050]

0070

[Chemical Formula 7]

□0051□

[0051]

[Chemical Formula 8]

□0052□

0000000

00001,3-00000000000000000000000002,

2-00000-1,3-0000000002-00000-2-000000-1,3-

000000002,2-000000-

1,3-0000000002,2-

00(00000000)-1,3-

0000000002,2-0000000-

1,3-00000000002,2-

[0052]

With this invention, it can also use diether compound component (C) as.

If it is a diether compound, there is not restriction. Desirable ones are diether which possesses branched aliphatic hydrocarbon and/or cycloaliphatic hydrocarbon group even amongthem.

Among these, it can use 1 and 3 -diether desirably, especially, 2 and 2-diisobutyl -1,3- dimethoxy propane, 2- isopropyl -2- isopentyl -1,3- dimethoxy propane, 2,2- dicyclohexyl -1,3- dimethoxy propane, 2,2- bis (cyclohexyl methyl) - 1 and 3 - dimethoxy propane, 2,2- diisopropyl -1,3- dimethoxy propane, 2-cyclopentyl -2- isopropyl -1,3- dimethoxy propane, 2-cyclopentyl -2- s-butyl -1,3- dimethoxy propane, 2-cyclohexyl -2- isopropyl -1,3- dimethoxy propane, 2-cyclohexyl -2- isopropyl -1,3- dimethoxy propane, 2-

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	cyclohexyl -2- s-butyl -1,3- dimethoxy propane, 2- isopropyl -2- s-butyl -1,3- dimethoxy propane, 2- phenyl -2- s-butyl -1,3- dimethoxy propane, 2- phenyl -2- s-butyl -1,3- dimethoxy propane, 2- benzyl -2- isopropyl -1,3- dimethoxy propane, 2- benzyl -2- s-butyl -1,3- dimethoxy propane, 2- (1 -methyl butyl) - 2 -isopropyl -1,3- dimethoxy propane, 2- (1 -methyl butyl) - can use 2 -s-butyl -1,3- dimethoxy propane desirably
DD(C)DDDDDDDDDDDDDDD(B) DDDDDDDDDDDDDDDDDDDD	amount used when component (C) is used, is inside range of 0.01 - 100 and inside range of preferably 0.1 ~ 10 , with mole ratio for component (B).
□0053□	[0053]
DOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOOO	With formation >this invention of <catalyst (a)="" (a),="" (b)="" according="" and="" as="" at="" be="" being="" both="" can="" catalyst="" coexisting="" component="" component,="" components="" consists="" crossing,="" dividing="" for="" form="" fourth="" inside="" into="" it="" itcontacts="" kind="" need="" of="" olefin="" one="" or="" outside="" polymerization="" polymerized,="" several="" something="" stepwise,="" td="" the="" the(c),="" this="" time,="" times="" to="" under="" underexisting="" vessel="" which="" with.<=""></catalyst>
□0054□	[0054]
DD(A)D(B)DDD(C)DDDDDDDDDDDDDDDDDDDDDDDDDDDDD	component (A), (B) or there is not especially restriction in supply method to contacting site of (C). It is normal respectively dispersing to hexane, heptane or other aliphatic hydrocarbon solvent, each one separatelyto add to polymerization vessel.
00(A)0000000000(B)00000 C)000000000000000000000000000000	component (A) component (B) or component (C) with can also add to the polymerization vessel separately with state of solid .
□0055□	[0055]
30000	[Working Example(s)]
00000000000000000000000000000000000000	In Working Example below, using Perkin Elmer make DSCtype 2, it did differential thermal analysis (DSC measurement)which measures melting point.
0001	Working Example 1

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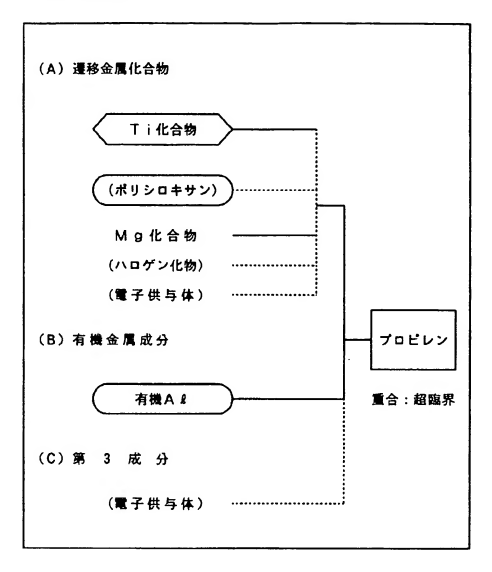
□□□(A)□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	In flask which nitrogen substitution is done, dehydration and 10 gram and and suspension state it introduced toluene 100ml which deoxygenation is done in {Production of component (A)} satisfactory, next Mg (OEt) ₂ introduced made.				
□□□□TiCl₄ 20□□□□□□□□□□□□□□□□-2- □□□□□□□□□□□□□□□□□□	Next, it introduced TiCl ₄ 20ml, temperature rise did in 90 deg C andintroduced 2 -isopropyl -2- isopentyl -1,3-dimethoxy propane 1.9ml, temperature rise did in 110 deg C and 3 hours reacted.				
00000000000000000	After reaction termination, you washed with toluene.				
□□□TiCl₄ 20□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	Next, it introduced TiCl ₄ 20ml and toluene 100ml, 2 hours reacted with 110 deg C.				
00000n- 00000000000000(A)0000 0000000(A-1)0000	After reaction termination, with n-heptane washing in satisfactory, solid component inorder to produce component (A) (A-1) with it did.				
	This titanium content was 2.5 weight %.				
□0056□	[0056]				
00000000000000000000000000000000000000	Next, in autoclave which nitrogen substitution is done, nheptane which wasrefined in same way as description above 200 ml was introducedin satisfactory, solid component which is synthesized at description above 5 gram was introduced, triethyl aluminum 1.5gram was introduced next, 4 - methylpentene -1 the prepolymerization was executed with 20 deg C.				
0000000n- 00000000000000(A)0000	After prepolymerization ending, with n-heptane washing in satisfactory, component (A) with it did.				
00000000000000000000000000000000000000	Removing portion, when you inspected amount of prepolymerization they were 10.6 gram poly 4-methylpentene -1/gram solid component (A-1).				
□□□□4-□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	Furthermore melting point of poly 4- methylpentene -1 was 231 deg C.				
□0057□	[0057]				
CO1000000000000000000000000000000000000	In stainless steel autoclave of internal volume 1.0liter which possesses {Polymerization of propylene } chuming and temperature control equipment in satisfactory dehydration and propylene which deoxygenation is done triethyl aluminum 8.9milligram was introduced with 25 deg C 100 ml, component (B) as.				
00000000000000000000000000000000000000	While introducing propylene into autoclave, temperature rise, increased pressure it did and itdesignated reaction system as supercritical state to 180 deg C, 100kg/cm ² .				

00000000000(A)000(A- 1)000100000000000000000000000000000000	Next, 10 milligram it introduced component (A) which is produced atdescription above with component (A-1) standard, started polymerization and 45 min polymerized.				
000000000000150000000	After polymerization termination, ethanol 15 ml was added, reaction was stopped.				
104	polymer of 104 gram was acquired, with MFR =51.8g/10 min, it was a I.I=95.3wt% with boiling n-heptane extraction.				
	In addition, Q was 6.2.				
□0058□	[0058]				
	Working Example 2				
	In flask which nitrogen substitution is done, dehydration and n-heptane which deoxygenation is done it introduced 200 ml in {Production of component (A)} satisfactory, Mg Cl ₂ 0.8 mole introduced 0.4 mole, Ti (O-nC ₄ H ₉) ₄ next, 2 hours reacted with 95 deg C.				
	After reaction termination, it lowered temperature to 40 deg C, 48 ml introduced methyl hydro polysiloxane (Those of 20 centistokes.) next, 3 hours reacted.				
0000000n-000000000	solid component which it forms was washed with n-heptane.				
□0059□	[0059]				
00000000000000000000000000000000000000	Next, in flask which nitrogen substitution is done, n-heptane which wasrefined in same way as description above 50 ml was introducedin satisfactory, solid component which is synthesized at description above 0.24 mole was introduced with Mg atom conversion.				
□□□n-□□□□25□□□□□□SiCl₄ 0.4□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	Next, mixing SiCl ₄ 0.4mole to n-heptane 25ml, it introduced to flask with30 deg C, 30 min, 3 hours reacted with 70 deg C.				
00000n-0000000000	After reaction termination, you washed with n-heptane.				
□□□n- □□□025□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□□	Next, mixing phthalic acid chloride 0.024mole to n-heptane 25ml, it introduced to flask with 70 deg C, 30 min, 1 hour reacted with 90 deg C.				
00000n-000000000	After reaction termination , you washed with n-heptane .				
00000n-000000000	After reaction termination , you washed with n-heptane .				
□□□□TiCl₄ 25□□□□□□□□□□□□100 deg C□3□□□□□□□□□	Next, introducing TiCl ₄ 25ml, 3 hours it reacted with 100 deg C.				
00000n- 00000000000000(A)0000 0000000(A-1)0000	After reaction termination, with n-heptane washing in satisfactory, solid component inorder to produce component (A)(A-1) with it did.				

00000000002.600%0000	This titanium content was 2.6 weight %.
□0060□	[0060]
00000000000000000000000000000000000000	Next, in autoclave which nitrogen substitution is done, nheptane which wasrefined in same way as description above 200 ml was introducedin satisfactory, solid component which is synthesized at description above 5 gram was introduced, triethyl aluminum 1.5gram was introduced next, 3 -methyl butene -1 the prepolymerization was executed with 10 deg C.
000000n- 000000000000(A)0000	After prepolymerization ending, with n-heptane washing in satisfactory, component (A) with it did.
00000000000000000000000000000000000000	Removing portion, when you inspected amount of prepolymerization they were 12.7 gram poly 3- methyl butene -1/gram solid component (A-1).
□□□□3-□□□□□□□□□302 deg C□□□□□	Furthermore melting point of poly 3- methyl butene -1 was 302 deg C.
□0061□	[0061]
0000000000000- 100000000000000000000000	propylene was introduced into autoclave in same way as {Polymerization of propylene } Working Example -1.
	Next (CH ₃) ₃ CSi (n-C ₃ H ₇) (OCH ₃) ₂ 1.0milligram, hydrogen 5 ml was introduced next triethyl aluminum 4.7milligram, component (C) as component (B) as.
	While introducing propylene into autoclave, it did temperature rise, increased pressure and itdesignated reaction system as supercritical state to 200 deg C, 150kg/cm ² .
00000000000(A)000(A- 1)00030000000000000045 0000000000000	Next 3 milligram it introduced component (A) which is produced atdescription above with component (A-1) standard and started thepolymerization and 45 min polymerized and stopped reaction in thesame way as Working Example -1.
127	polymer of 127 gram was acquired, with MFR =68.2g/10 min, it was a I.I=97.1wt% with boiling n-heptane extraction.
000Q007.100000	In addition, Q was 7.1.
□0062□	[0062]
□□□3~6	Working Example 3~6
000-20000000(A)000000- 100000(B)0000(C)00000001 00000000000000000200	Using component (A) which is produced with Working Example -2, component whichit shows in Table 1 (B) and using component (C), other than making the polymerization condition which it shows in Table 1, you polymerized with condition which is completely similar to Working Example 2.

	表 1										
	et/\ (D)	÷4. (0)	单量体成分		重合時		重合結果				
実施例	英施例 成分(B) 成分(C) (使用量)	臨界温度	庭界圧力	温 度 (℃)	圧 カ (kg/cn²)	\$97-収量 (g)	MER (g/105))	I. I (wt%)	QM		
実施例 1	ትባ፤ ተ ልፖል ፯ ፡፡ ሳል (8. 9mg)	-			180	100	104	51, 8	95. 3	6. 2	
実施例 2	ትሀ1 ታ ል7ቆ2፡፡ሳል (4. 7ng)	(CH ₃) ₃ CSi(N-C ₃ H ₇ OCH ₃) ₂ (lmg)				200	150	127	68, 2	97. 1	7. 1
実施例 3	· トワインフチルアルミニウム	CH ₈ CH ₈ H ₈ C - C CH ₃ CH ₃ / OCH ₈ CH ₃ CH ₃ / OCH ₈	70€VV 91. 4°C	foedy 45. 4kg/cm²	190	135	125	51. 6	96.8	7.3	
	(12mg)	(1.3mg)			į						
実施例 4	}9n-4\$%76\$29& (11.5mg)	t-C ₄ H ₉ Si OCH ₃ n-C ₄ H ₉ Si OCH ₈ (1. lng)			185	140	129	34. 7	97. 3	7, 6	
実施例		(C ₂ H ₅) ₂ Si(OCH ₈) ₂									
5	(18.5mg)	(1.9mg)			200	140	115	88. 3	95. 4	7.0	
実施例	トリローデジルブルミニウム	t-C ₄ H _B Si C ₂ H ₅ OC ₂ H ₅			210	170	111	117. 2	95. 2	6.8	
6	(27.3mg)	(2.5mg)									

	[Brief Explanation of the Drawing(s)]
0000	[Figure 1]
00000000000000000000000000000000000000	It is a flowchart in order to help understanding of this invention .
Drawings	
0000	[Figure 1]



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